# Package 'a4Core'

October 11, 2016

Type Package
Title Automated Affymetrix Array Analysis Core Package
<b>Version</b> 1.20.0
Date 2011-06-04
Author Willem Talloen, Tobias Verbeke
<b>Maintainer</b> Tobias Verbeke <tobias.verbeke@openanalytics.eu>, Willem Ligtenberg <willem.ligtenberg@openanalytics.eu></willem.ligtenberg@openanalytics.eu></tobias.verbeke@openanalytics.eu>
Description Automated Affymetrix Array Analysis Core Package
Depends methods, Biobase, glmnet
License GPL-3
biocViews Microarray
NeedsCompilation no

# **R** topics documented:

1		
simulateData	 	 2

confusionMatrix	Generic function to produce a confusion matrix (related to a classifi-
	cation problem)

# Description

Generic function to produce a confusion matrix (related to a classification problem)

#### Usage

confusionMatrix(x, ...)

#### Arguments

Х	object (usually a model fit object) that contains all information needed to pro- duce the confusion matrix.
	further arguments for a specific method

# Author(s)

Tobias Verbeke

simulateData Simulate Data for Package Testing and Demonstration Purposes

# Description

Simulate Data for Package Testing and Demonstration Purposes

# Usage

```
simulateData(nCols = 40, nRows = 1000, nEffectRows = 5, nNoEffectCols = 5,
betweenClassDifference = 1, withinClassSd = 0.5)
```

#### Arguments

nCols	number of samples; currently this should be an even number
nRows	number of features (genes)
nEffectRows	number of differentially expressed features
nNoEffectCols	number of samples for which the profile of a differentially expressed feature will be set similar to the other class
betweenClassDifference	
	Average mean difference between the two classes to simulate a certain signal in the features for which an effect was introduced; the default is set to 1
withinClassSd	Within class standard deviation used to add a certain noise level to the features for which an effect was introduced; the default standard deviation is set to 0.5

# Value

object of class ExpressionSet with the characteristics specified

#### Note

The simulation assumes the variances are equal between the two classes. Heterogeneity could easily be introduced in the simulation if this would be requested by the users.

# topTable

# Author(s)

W. Talloen and T. Verbeke

# Examples

```
someEset <- simulateData(nCols = 40, nRows = 1000, nEffectRows = 5, nNoEffectCols = 5)
someEset</pre>
```

topTable

S4 Generic for obtaining a top table

# Description

a top table is a rectangular object (e.g. data frame) which lists the top n most relevant variables

# Usage

topTable(fit, n, ...)

#### Arguments

fit	object for which to obtain a top table, generally a fit object for a given model class
n	number of features (variables) to list in the top table, ranked by importance
	further arguments for specific methods

# Author(s)

Tobias Verbeke

topTable-methods *Methods for topTable* 

# Description

Methods for topTable. topTable extracts the top n most important features for a given classification or regression procedure

# Arguments

fit	object resulting from a classification or regression procedure
n	number of features that one wants to extract from a table that ranks all features according to their importance in the classification or regression model; defaults to 10 for limma objects

# Methods

glmnet and lognet

glmnet objects are produced by lassoClass (a4Classif) or lassoReg (a4Base)

# Index

\*Topic manip simulateData, 2 topTable-methods, 3 \*Topic methods topTable-methods, 3 \*Topic models confusionMatrix, 1 topTable, 3

confusionMatrix, 1

simulateData, 2

topTable, 3
topTable,elnet-method
 (topTable-methods), 3
topTable,glmnet-method
 (topTable-methods), 3
topTable,lognet-method
 (topTable-methods), 3
topTable-methods, 3